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- (54)METHOD FOR LOWERING POLLEN FERTILITY BY USING POLLEN-SPECIFIC ZINC FINGER TRANSCRIPTIONAL FACTOR GENES
- A method is provided for producing a male sterile plant by utilizing a plant expression cassette including a nuclaic acid which is DNA ancoding zinc finger transcription factors (ZPT2-5, ZPT3-1 and ZPT4-1) derived from Patunia and a promoter oparatively linked to the nucleic acld. Further, a method is provided for producing

a plant having a modified trait by utilizing a plant exprassion cassette including a promoter derived from the ZPT3-1 and ZPT4-1 genas and a hetarologous gene operatively linked to the promoter.

### Description

### TECHNICAL FIELD

[0001] The present invention relates to genes which are expressed specifically in the pollen producing its-sues of stamens and use of the same. More particularly, the present invention relates to the genes for zinc finger imascription factors (ZPT2-5, ZPT3-1 and ZPT4-1) derived from Petunia, which are expressed specifically in ...10 microspores, and use of the same.

#### BACKGROUND ART

[0002] Pollen fertility causes problems in various aspects of agriculture and horticulture. For example, in the case of mating for cross breeding, self-pollination has to be avoided by castration (removal of stamens) which requires enormous effort. In the seed and seedling industry, there is a demand for a trait of lack of pollen fer- 20 tility from the standpoint of commercially protecting excellent breeds obtained by cross breeding. To meet such a demand, a technique for controlling poilen fertility (pollination control) has been strongly required. Conventionally, for particular crops, lines of cytoplasmic male sterlity have been used for cross breeding, and some success has been achieved. However, the cytoplasmic sterility trait is often accompanied by undesired side effects, such as a reduction in disease resistance and the like. There are further problems, such as that the trait is unstable, that it is difficult to mass-produce the seeds. and the like. A method for reducing the fertility by treating with a chemical agent(s) has been studied, but safety evaluation and elucidation of the mechanism of this method have not been fully done and thus such a method is not yet in actual use. Therefore, there is a demand for an excellent male sterilization technique using genetic engineering.

[0003] Pollen is the male gametophyte of spermatophyte. The development of pollen which proceeds while 40 pollen is surrounded by an anther as a supporting tissue is divided into the following stages: the tetrad stage immediately after the meiosis of microsporogenous cells (pollen mother cells); the release stage during which microspores are released from the tetrad; the uninucleate 45 stage characterized by the enlargement and vacuoiation of pollen cells, the mitotic stage giving rise to the differentiation into vegetative and generative cells by mitosis; and the subsequent binucleate stage. After these stages, the anther finally dehisces and matured pollen 50 grains are released. Therefore, it can be said that the microspore is one of target tissues which are most suitable for artificial control in order to inhibit the development of polien and eliminates pollen fertility.

[0004] As described above, great expectations are placed on male sterilization techniques using genetic engineering. Particularly, if a gene which is expressed specifically in the direct precursor of a pollen cell, such

as a microspore, can be utilized. It is considered to be highly likely that male sterilization can be achieved without conferring undesired traits to plants. Several examples of promoters specific to various stamen tissues and gene constructs for male sterilization comprising the promoter have been reported (Shivanna and Sawhney Ed., Pollen biotechnology for crop production and improvement (Cambridge University Press), pp. 237-257, 1997). However, there has been continuously a demand for a novel gene useful for control of pollen fertility, which has high tissue and temporal specificities of expression. [0005] Recently, the inventors of the present application specified the cDNA sequences of novel transcription factors derived from Petunia, i.e., seven zinc finger (ZF) transcription factors including PEThy ZPT2-5. PEThy ZPT3-1, and PEThy ZPT4-1 (hereinafter abbreviated as ZPT2-5, ZPT3-1, and ZPT4-1, respectively). And the inventors reported that Northern blot analysis indicates that each transcription factor transiently expresses in an anther-specific manner in a different stage of the development of the anther (Kobayashi et al., Plant J., 13:571, 1998). However, the physiological function and action of these transcription factors in plants, and the precise expression sites and the expression controlling mechanism of the genes encoding the transcription factors have been not clarified.

### DISCLOSURE OF THE INVENTION

[0006] The objective of the present invention is to provide a genetic engineering technique using a pollenspecific gene which is useful for modification of a plant trait, representatively male sterility.

[0007] The present inventors raintroduced genes encoding anther-specific transcription factor (ZPT2-5,
ZPT3-1 and ZPT4-1), which had been previously isolated from Petuna, into Petunia. As a result, it was found
that the normal development of polien was inhibited, so
that polien fertility was significantly reduced (ZPT2-5
de and ZPT4-1), or substantially eliminated (ZPT3-1). Further, the inventors isolated upstream regions of the
ZPT3-1 and ZPT4-1 genomic genes, respectively, and
studied the tissue specificity of the promoter activity, As
a result, it was found that the promoter activity is exds pressed in microspores from the uninucleate stage to
the binucleate stage in a tissue and temporal-specific
manner. The present invention was completed based on
these findings.

[0008] According to a first aspect of the present invenor tion, a method for producing a male sterile plant comprises the steps of providing a plant expression cassette including: a nucleic acid being any of () DNA having a lasequence from position 1 to position 777 of a base sequence indicated by SEO ID NOT: 1, (ii) DNA hyvindzing to the DNA having the base sequence (i) under stringent of conditions and encoding a transcription factor controlling the development of pollen, and (iii) a DNA fragment of (ii) or (ii); and a promoter operatively linked to the nucleic soid, providing plant cells having an endogenous transcription factor controlling the development of polien, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions, introducing the expression cassette into the plant cells, regenerating the plant cells, into which the expression cassette has been introduced, to plants; and screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppression.

[0009] According to a second aspect of the present invention, a method for producing a male sterile plant comprises the steps of providing a plant expression cassette including: a nucleic acid being any of (i') DNA hav-Ing a sequence from position 1 to position 1640 of a base sequence indicated by SEQ ID NO: 3, (iii) DNA hybrid-Izing the DNA having the base sequence (i') under stringent conditions and encoding a transcription factor controlling the development of pollen, and (iii') a DNA fragment of (i') or (ii'); and a promoter operatively linked to the nucleic acid, providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions, introducing the expression cassette into the plant cells, regenerating the plant cells. into which the expression cassette has been introduced. to plants; and screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed. [0010] According to a third aspect of the present invention, a method for producing a male sterile plant comprises the steps of providing a plant expression cassette including: a nucleic acid being any of (I") DNA having a sequence from position 1 to position 1948 of a base sequence indicated by SEQ ID NO: 5, (ii\*) DNA hybridizing the DNA having the base sequence (I") under stringent conditions and encoding a transcription factor controlling the development of pollen, and (III") a DNA fragment of (I") or (II"); and a promoter operatively linked to the nucleic acid, providing plant cells having an endogenous transcription factor controlling the development of polien, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions, introducing the expression cassette into the plant cells, regenerating the plant cells, into which the expression cassette has been introduced, to plants; and screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed. [0011] It should be noted that the DNAs of (ii), (ii') and (ii") each do not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of abase sequence indicated by SEQ ID NO: 13 under stringent conditions and encoding a transcription factor controlling the development of pollen. The base sequence indicated by SEQ ID NO: 13 is a cDNA sequence encoding another transcription factor ZPT3-2 isolated from

Petunia (Kobayashi et al. above).

[0012] The method according to the first through third aspects of the present invention is utilized as a method for conferring male sterility to a plant.

- [0013] In one embodiment of the first through third aspects, the nucleic acid is linked in a forward direction with respect to the promoter, and may be transcribed in a sense direction in cells of the plant.
- [0014] In one embodiment of the first through third aso pects, the nucleic acid is linked in a reverse direction with respect to the promoter, and may be transcribed in a antisense direction in cells of the plant.
- [0015] In one embodiment of the first through third aspects, the plant is dicotyledon. The dicotyledon is preferably of the family Solanaceae, and more preferably of the genus Petunia.
- [0016] In one embodiment of the first through third aspects, the expression cassette is incorporated into a plant expression vector.
- 20 [0017] According to the first through third aspects of the present invention, a male sterile plant produced by a method according to any of the above-described methods is also provided.
- [0018] According to a fourth aspect of the present invention, a method for producing a plant having a modified trait comprises the steps of providing a plant expression cassette including: a promoter including any of (a')
  DNA having a sequence from position 1 to position 2824
  of a base sequence indicated by SEQ ID NO. 7 and (b')
  DNA having a part of the sequence of (a') and exhibiting
  promoter activity specific to microspores; and a heterologous gene operatively linked to the promoter, introducing the expression cassets that plant cells, and regenerating the plant cells, into which the expression cassettle has been introduced, to plants.
- [0019] According to a fifth aspect of the present invention, a method for producing a plant having a modified trall comprises the steps of providing a plant expression cassette including; a promoter including any of (a") DNA 40 having a sequence from position 1 to position SS1 of ra bases sequence inclusated by SEQ ID NO: 8 and (b") DNA having a part of the sequence of (a") and exhibiting promoter activity specific to microspores and optionally the chisocone tissue of an arther; and a heterologous
- motor activity specific to microspores and optionally the dehisconce tissue of an anther, and a heterologous 5 gene operatively linked to the promoter, introducing the expression cassette into plant cells, and regenerating the plant cells, into which the expression cassette has been introduced, to plants.
- [0020] In one embodiment of the fourth and fifth aspects, the trait is fertility, and the plant having a modified trait is a male sterile plant. Therefore, the method of the present invention may be utilized as a method for conferring male sterlity to a plant.
- [0021] In one embodiment of the fourth and fifth aspects, the trait is compatibility, and the plant having a modified trait is a self-incompatibile plant. Therefore, the method of the present invention may be utilized as a method for conferring self-incompatibility to a plant.

[0022] In one embodiment of the fourth and fifth aspects, the plant is dicotyledon. The dicotyledon is preferably of the family Solanaceae, and more preferably of the genus Petunia.

[0023] In one embodiment of the fourth and fifth aspects, the expression cassette is incorporated into a plant expression vector

[0024] In one embodiment of the fourth and fifth aspects, a trait-modified plant produced by a method according to any of the above-described methods is pro-

[0025] According to a sixth aspact of the present invention, a promoter comprises DNA of the following (f) or (II): (f) DNA having a sequence from position 1 to poelltion 2624 of a base sequence Indicated by SEO ID 19 NO: 7, and (II) DNA having a part of the sequence (f) and exhibiting promoter activity specific to microsporces.

[0036] According to a seventh aspect of the present invention, a promoter comprises DNA of the following 20 (1) or (III); (II) DNA having a sequence from position 1 to position 938 of a base sequence indicated by SEO ID NO: 8; and (III) DNA having a part of the sequence of (II) and exhibiting promoter activity specific to microspores and optionally the dehiscence tissue of an anther.

10027] According to an eighth aspect of the present invention, a plant expression casseste useful for confering male stellity to a plant, comprising any of the above-described microspore-specific promoters and a pheterologous gene operatively linked to the promoter.

BRIEF DESCRIPTION OF THE DRAWINGS

### [0028]

Figure 1 is a diagram showing a cDNA sequence of a gene encoding ZPT2-5 (herein also simply referred to a "ZPT2-5 gene") and the corresponding amino acid sequence. Two zinc finger motifs and a 40 DLNL sequence (amino acids from position 145 to position 1450 are underline).

Figure 2 is a diagram showing a cDNA sequence of a gene encoding ZPT3-1 (herein also simply referred to as "ZPT3-1 gene") and the corresponding amino acid sequence. Three zinc finger motifs and a DLNL sequence (amino acids from position 408 to position 417) are underlined.

Figure 3 is a diagram showing a cDNA sequence of a gene encoding ZPT4-1 (herein also simply referred to as "ZPT4-1 gene") and the corresponding amino acid sequence. Four zinc finger motifs and a DLNL sequence (amino acids from position 438 to 55 position 449 are underlined.

Figure 4 is a schematic diagram showing structures

of plant expression vectors used for expression of each cDNA sequence of ZPT2-5, ZPT3-1 and ZPT4-1 (pBIN-35S-ZPT2-5, pBIN-35S-ZPT3-1 and pBIN-35S-ZPT4-1).

Figure 5 is a diagram showing an upstream sequence of the coding region of the ZPT3-1 gene. The transcription initiation site is indicated by a thick arrow (position 2567). The translation initiating codon (ATG) is indicated by a thick underline.

Figure 6 is a diagram showing an upstream sequence of the coding region of the ZPT4-1 gene. The transcription initiation site is indicated by a thick arrow (position 3503). The translation initiating codon (ATG) is indicated by a thick underline.

Figure 7 is a schematic diagram showing structures of plant expression vectors for analyzing promoters for the ZPT3-1 and ZPT4-1 genes (pBIN-ZPT3-1-GUS),

Figure 8 shows photographs indicating the forms of organisms, i.e., the pollen of a wild type Petunia and the pollen of a Petunia into which pBIN-35S.ZPT2-5 was introduced (a transformatin in which cosupression occurred) (the magnification is 400 times). Figures 8(a) through (d) are of the wild-type Petunia and Figures 8(e) through (f), are of the cosupressed transformed Petunia, each of which shows the polen of a bud at a different development stage. All the pollen were stained by a commonly each wed method using DAPI (4', 8-diamidino-2-phenylindole dihydrochloride n-hydrate).

Figure 9 shows photographs indicating the forms of organisms, Is, in the polien of a walf type Petunia and the polien of a Petunia into which pBIN-35S-ZFT3-1 was introduced (the magnification is 700 times). Figures 9(a) and (e) are of the wild-type Petunia and Figures 9(b) and (e) are of the wild-type Petunia and Figures 9(b) and (e) are of the renatformed Petunia, each of which shows the polien at the tetrad stage and the microspore stage, respectively. The polien of the tetrad stage and the polien of the microspore stage were stained by a commonly used method using DAPI and safranin, respectively. The polien of the Petunia into which pBIN-35S-ZFT4-1 was produced showed substantially the same form as Figures 9(b) and 9(c).

Figure 10 shows photographs showing the forms of organisms, i.e., GUS-statined floral organs of Petunia into which pBIN-2713-1-GUS and pBIN-2714-1-GUS were introduced. Each photograph was taken of a llower (bud) whose anther is in the unlinucleate stage. Figures 10(a) and (d) show the appearances of bud at the actual size. Figures 10(b) and (d) show the cross-sectional views

of an anther at a low magnification (40 times). Figures 10(c) and (f) show the cross-sectional views of microspores (Figure 10(c); the magnification is 700 times) and the dehiscence tissues and the surrounding vicinity of the anther (Figure 10 (f); the magnification is 200 times) at high magnifications.

BEST MODE FOR CARRYING OUT THE INVENTION

[0029] Hereinafter, the present invention will be described in detail.

(Transcription factors derived from ZPT2-5, ZPT3-1 and ZPT4-1 genes)

[0030] A nucleic acid, which is useful in a method for producing male sterile plants according to first to third aspects of the present invention, is any one of the following DNAs:

- (i) DNA having a sequence from position 1 to position 777 of a base sequence indicated by SEQ ID
- (ii) DNA having a sequence from position 1 to position 1640 of a base sequence indicated by SEQ ID NO: 3;
- (I") DNA having a sequence from position 1 to position 1948 of a base sequence indicated by SEQ ID NO: 5.

[0031] DNA which hybridizes to the DNA having any of the base sequences (i) to (I") under stringent conditions, and encodes a transcription factor which controls the development of polien(i.e., (ii), (ii') or (ii")); or DNA which is a fragment of any of the above-de-

scribed DNAs (i.e., (iii), (iii') or (iii")).

[0032] The above-described nucleic acid of the present invention is preferably DNA of (i), (i') or (i"), i.e., DNA encoding ZPT2-5, ZPT3-1 or ZPT4-1, or a fragment thereof, and more preferably DNA of (i), (i') or (i"). [0033] in the present specification, "transcription factor" refers to a protein for controlling the synthesis of mRNA by binding to DNA in the regulatory region of a gene. It is known that a certain type of transcription factor has a highly conserved amino acid sequence called a zinc finger (ZF) motif in the DNA binding domain. ZPT2-5 is a zinc finger (ZF) protein of the Cvs2/His2 type (EPF family), which is a transcription factor which includes two ZF motifs in the full-length amino acid sequence consisting of 176 amino acids, and further, a hydrophobic region called a DLNL sequence. Similarly, ZPT3-1 is a ZF protein of the EPF family, which is a tran-

scription factor which includes three ZF motifs in the full-

length amino acid sequence consisting of 437 amino ac-

a ZF protein of the EPF family, which is a transcription

factor which includes four ZF motifs in the full-length

amino acid sequence consisting of 474 amino acids.

and further, a DLNL sequence. For any of the abovedescribed transcription factors, see Kobayashi et al. (above). cDNA sequences (SEQ ID NO: 1, 3 and 5) encoding ZPT2-5, ZPT3-1 and ZPT4-1, respectively, are shown in Figures 1, 2 and 3 along with corresponding putative amino acid sequences (SEQ ID NO: 2, 4 and 6). [0034] In the present specification, "fragment" of a nucleic acid or DNA refers to a fragment which can inhibit the expression of an endogenous transcription factor in a plant when the fragment is introduced into the plant and expressed in an appropriate manner. This fragment is selected from regions of DNAs of the above-described (i), (i'), (l''), (li), (li') or (it") other than the regions encoding the zinc finger motifs in the DNAs. The fragment has a length of at least about 40 bases or more, preferably about 50 bases or more, more preferably about 70 bases or more, and even more preferably about 100 bases

[0035] In the present specification, "stringent conditions" for hybridization are intended as conditions sufficient for the formation of a double-strand oligonucleotide of a particular base sequence (e.g., DNA encoding ZPT2-5, ZPT3-1 or ZPT4-1 derived from Petunia) and another base sequence having a high level of homology with the particular base sequence (e.g., DNA encoding a homolog of ZPT2-5, ZPT3-1 or ZPT4-1 which is present in a plant other than Petunia). A representative example of the stringent conditions applied to the present invention are the following: hybridization is conducted in a solution containing 1M NaCi, 1%SDS, 10% dextran sulfate, 32P-labeled probe DNA (1×107 cpm) and 50 µg/ml salmon sperm DNA at 60°C for 16 hours. followed by washing twice with 2×SSC/1%SDS at 60°C for 30 minutes.

[0036] In the present invention, a degenerate primer pair corresponding to a conserved region of an amino acld sequence encoded by the gene of a known transcription factor may be used in order to isolate DNAs encoding ZPT2-5, ZPT3-1 and ZPT4-1, and DNA encoding a transcription factor which hybridizes these DNAs under stringent conditions to inhibit the development of pollen. PCR is conducted using this primer pair with cDNA or genomic DNA of a plant as a template, thereafter, the resultant amplified DNA fragment is used as a probe so that the cDNA or genomic library of the same plant can be screened. As an example of such a primer pair, a combination of 5'-CARGC-NYTNGGNGGNCAY-3' (SEQ ID NO: 9), and 3.-RT-GNCCNCCNARNGCYTG-5' (SEQ ID NO: 10) is illustrated (where N indicates inosine, R indicates G or A, and Y indicates C or T). The above-described primer sequences each correspond to an amino acid sequence QALGGH included in the zinc finger motifs of the abovedescribed ZPT transcription factors.

ids, and further, a DLNL sequence. Similarly, ZPT4-1 is 55 [0037] Therefore, the stringent hybridization conditions which are applied to the present invention may also be used for PCR. In a representative example, the above-described degenerate primers (SEQ ID NOs: 9 and 10) may be used. In this case, the PCR reaction conditions may be the following: denaturation at 94°C for 5 minutes; followed by 30 cycles of 30 seconds at 94°C, 30 seconds at 50°C and 1 minute at 72°C; and finally, incubation at 72°C for 7 minutes.

[0038] PCR may be conducted based on the manufacturer's instruction for a commerciality available kit and device, or a method well known to those skilled in the art. A method for preparing a gene library, a method for cloning a gene, and the like are also well known to those skilled in the art. For example, see Sambrook et al., Mociecular Cloning: A Laboratory Manual, 2nd Ed. (Cold Spring Flarbor Laboratory, 1989). The base sequence of a resultant gane may be determined with a nucleotide saquencing analysis mathod known in the art, or by a commercially available automatic sequencer.

[0039] In the prasent specification, "controlling the development of polian" by a transcription factor representatively means that when the expression of this transcription factor is inhibited, a significant change in the form or functions of polien is observed. Representatively, by inhibiting the axpression of a gene encoding the transcription factor of the present invention, preferably about 75% or more, more preferably about 95% or more of polien colls are killed before being matured. When the amount of mRNA measured by a Northern blot method is about one tenth or ioss as compend to a wild-type control plant, the expression of a transcription factor is judged to be inhibited.

10040) Whether or not the transcription factors encoded by genes located and identified by screening as above (i.e., ZPT2-5, ZPT3-f and ZPT4-f, and the homologs thereof) control the development of pollen, can be confirmed by producing a transformed plant and observing the characteristics of the pollen of the plant in accordance with the disclosure of the present specifica-

[0041] According to the present invention, DNA enodding at reascription factor which controls the development of pollen can be utilized to inhibit the expression of an endogenous gene having the same or homologous base sequence as that of the DNA in plant cells. Such a target endogenous gene is also a transcription factor which controls the development of pollen. According to the method of the present invention, plants are conferred mails sterility by selectively inhibiting only the expression of an endogenous transcription factor, prefersibly without substantially inhibiting the expression of genes other than the endoganous transcription factor which controls the development of pollen.

[0042] In other words, plant cells to which the expression inhibiting lechnique of the present invention is applied are plant cells having an endogenous transcription factor which controls the development of pollen. The gene encoding this endogenous transcription factor is defined as a gene which hybridizes with DNA encoding the above-described ZPT2-5, ZPT3-1 or ZPT4-1, or a homolog thereof under stringent conditions. The dafinition of the "stringent conditions" is the same as that described in relation to specification of the homologs of ZPT2-5, ZPT3-1 and ZPT4-1. Plants capable of being conferred male sterility with the above-described method are preferably plants which are phylogenetically, closely related to Petunia from which the above-described ZPT genes are isolated, or plants from which genes encoding the above-described ZPT homologs are isolated, but the present invention is not intended to be limited to this. "Plants which are phylogenetically, closely related" means representatively plants categorized into the same order, preferably categorized into the same family, more preferably categorized into the same genus, and even more preferably categorized into the same species. Considering the fact that the development of pollen is essantial for the reproduction of spermatophyte, it could be easily understood that transcrip-

tion factors having the same or similar function to that

of ZPT2-5, ZPT3-1 and ZPT4-1 may be widely present

in other plants.

[0043] As a technique for suppressing the expression of an endogenous gene, cosuppression and antisense techniques may be utilized, representatively. As to cosuppression, when a recombinant gene is introduced into a plant cell, the expression of both the gene Itself and an endogenous gene including a sequence homologous to part of that gene are suppressad. When cosupprassion is utilized, an expression cassette according to the present invention includes DNA encoding a transcription factor or a fragment thereof in the form linked in a forward direction with respect to the promoter. After DNA encoding a transcription factor or a fragment thareof is introduced into a plant ceil as an expression cassette, the DNA or fragment thereof can be transcribed in the sense direction under control of the promoter. Due to the action of the introduced DNA, it is possible to suppress the targeted gene expression. Cosuppression can be observed in some transformed plant individuals, but mostly, cosuppression does not occur sufficiently in other individuals. Therefore, typically, individuals in which gene expression is suppressed in an intended manner are screened with routine procedures.

[0044] Antisense means that when a recombinant 45 gene is introduced into a plant cell, the transcribed product (mRNA) of the introduced gene forms a hybrid with the complementary sequence of the transcribed product (mRNA) of an endogenous gene so that the translation of a protien encoded by the endogenous gene is inhibited. When antisense is utilized, the expression cassette of the present invention includes DNA encoding a transcription factor or a fragment thereof in the form linked in a reverse direction with respect to the promoter. After DNA encoding a transcription factor or a fragment thereof so its introduced into a plant cell as an expression cassette, the DNA or fragment thereof may be transcribed in the antisense direction under control of the promoter. Due to the action of the entisense transcripts, it is pos-

sibla to suppress the expression of the targeted gene.

(Promotars derivad from ZPT3-1 and ZPT4-1 genes)

[0045] A promoter useful in a method for producing a plant having a modified trait according to the fourth and fifth aspects of the Invention is a promoter which Includes any of (a') DNA having a sequence from position 1 to position 2624 of a base sequence indicated by SEQ ID NO: 7; (a") DNA having a sequence from position 1 to position 3631 of a base sequence Indicated by SEQ ID NO: 8; and DNA having a part of tha sequences (a') or (a") and which exhibits promoter activity specific to microspores. The above-described promoter of tha presant Invantion is praferably the promoter of (a') or (a"), i.e., the promoter for the ZPT3-1 or ZPT4-1 gene. [0046] A sequence having promoter activity specific to microsporas, which is obtained by removing a sequance which is not essantial for tissua-specific axprassion activity from the promotar regions for the ZPT3-1 and ZPT4-1 genes, falls within the scope of the present invention. Such a saquanca can be obtained by conducting a promoter dalation experiment in accordance with a commonly used method, Briefly, a plasmid obtained by fusing various promoter ragion delation mutants of the ZPT3-1 or ZPT4-1 gene (e.g., mutants obtained by deleting the promoter region from the 5' upstraam side of the ZPT3-1 or ZPT4-1 gana in various lengths), and an appropriata reporter gene (e.g., the GUS gene) can be used to measure the tissue-specific 30 promotar activity of the deletion mutants, thereby identifying a ragion essential for the activity.

[0047] Onca the region essential for the promoter activity is identified, it is possible that a sequence within or adjacent to the region is modified so that the magnitude of the expression activity of the promoter is increased. The thus-obtained variants also fall within the present invention as long as the variants exhibit promoter activity spacific to microspores.

[0048] In the present Invention, "exhibit promoter activity specific to microspores" means that the ability of a promoter to Initiata the transcription of DNA to direct gene expression in a naturally-occurring plant or a plant to which the promoter is introduced as an expression cassatta in which the promoter is linked to an arbitrary structural gene, is exhibited specifically in microspores. Hare, "specific" means that the expression activity of a promoter is higher than in all tha other tissues of the flower of the same plant (including tapetum layer, filament, style, capitulum, petal, calyx, and the like; note that the dehiscence tissue of the anther is excluded). The above-described specific promoter preferably has an exprassion activity in microspores, higher than the expression activity in all the other tissues of the flower and portions other than the flower of the same plant 55 (roots, laavas, stams, and tha like). More preferably, the specific promoter exhibits substantially no activity in all the other tissues of the flower and portions other than

the flower of the same plant. "Exhibit the promoter activity specific to the dehiscence tissue of the anther" is defined in a manner similar to that described above. The magnitude of expression activity may be evaluated by comparing the expression level of a promoter in microspores with the expression level of the same promoter in other flower tissues in accordance with a commonly used method. The expression level of a promoter is typically datarmined by the production amount of the products of a gene expressed under control of the promoter. [0049] The above-described method of the present invention utilizing a specific promoter is intended to modify a trait related to reproduction of a plant. "Modify" means that at least a portion of the reproductive organ of a posttransformation plant loses a function which existed in the pre-transformation plant (wild type or horticulture bread), acquires a function which did not exist in the pretransformation plant, or has an increased or decreased level of particular function as compared to the pre-transformation plant. Such modification of a trait can be achieved as a result of the microspore-specific expression of any hetarologous gane operatively linked to the promotar of the presant invention under the control of the promoter in a transformed plant into which the gene has been introduced. It is well known that in a number of tissue-specific promoters, tha tissue-specificity is conserved among species. Therefore, it is easily undarstood that the promoter of the present Invention can be applied to a wide variety of plant species. The degree of trait modification may be evaluated by comparing the trait of a post-transformation plant with the trait of the pre-transformation plant. As a preferable trait to be mod-Ified, female sterility and self-incompatibility are illustrated, but such a trait is not limited to thesa.

[0050] For axampla, the promoter of the present invention can be obtained by screening the genomic library of a plant using known CDNA as a probe, and declaring an upstream sequence of a coding region from the corrasponding genomic clona. As an axampla of cD-NA, CDNA of the above-described transcription factors derived from Patunia, 2PT3-1 and ZPT4-1, ser illustrations.

[0051] The promoter of the present Invention is not imited to that lealeted from the nature, but may include imited to that lealeted from the nature, but may include youtheasted polynucleotides may be obtained by synthesizing or modifying the sequence of a promoter sequenced as described above or an active region thereof with a method well-known to those skilled in the art.

(Construction of expression cassette and expression vector)

[0052] DNA encoding the transcription factor of the ip present invention can be introduced into plant cells as an expression cassette, in which the DNA is oparatively linked to an appropriate promoter using a method well known to those skilled in the art, with a known gene recombinant technique. Similarly, the microspore-specific promoter of the present invention cen be introduced into plent cells as an expression cessette in which the promoter is operatively linked to a desired heterologous gene.

[0053] A "promoter" which can be linked to the abovedescribed transcription factor means any promoter which expresses in plants, including any of a constitutive promoter, a tissue-specific promoter, end en inducible promoter.

[0054] "Constitutive promoter" refers to a promoter which causes a structural gene to be expressed at a cartain level irrespective of stimuli inside or outside plant cells. When a heterologous gene is expressed in other tissues or orgens of a plant and a plant is not given an undesired trait, use of a constitutive promoter its simple and prefereble. As examples of such a constitutive promoter, as examples of such a constitutive promoter, as promoter (PSSS) of cauliflower mosalc virus (CAMV), and the promoter for nopaline synthase (Tnos) are illustrated, but the constitutive promoter is not limited to these.

[0055] In the present invention, "tissue-specific promoter" refers to a promoter which causes a structure! gene to be expressed specifically in at least microspores. Such a tissue-specific promoter includes the promoters derived from ZPT3-1 and ZPT4-1 genes of the present invention and, in addition, other known promoters having anther-specific expression activity. Therefore, use of an expression cassette of the naturally-occurring ZPT3-1 and ZPT4-1 genes comprising a microspore-specific promoter and a sequence encoding a transcription factor optionally combined with another regulatory element, falls within the present invention. [0056] "Inducible promoter" refers to a promoter which causes a structural gene to be expressed in the presence of a particular stimulus, such as chemical agents, physical stress, and the like, and which does not exhibit expression activity in the absence of the stimulus. As an example of such en inducible promoter, e glutathione S-transferase (GST) promoter which can be induced by auxin (van der Kop, D. A. et al., Plant Mol. Biol., 39:979, 1999) is illustrated, but the Inducible promoter is not limited to this.

[0057] In the present specification, the term "expression cassetie" or "plant expression cassetier "refers to a nucleic acid sequence including DNA encoding the transcription factor of the present invention and a plant expression promoter operatively (i.e., in such a manner that can control the expression of the DNA) inked to the DNA, and a nucleic acid sequence including the microspore-specific promoter of the present invention and a heterologous gene operatively (i.e., in-frame) linked to the promoter.

[0058] "Heterologous gene" which mey be linked to the ebove-described microspore-specific promoter refers to any of endogenous genes of Petunia other than the ZPT3-1 and ZPT4-1 gene, endogenous genes in a plant other then Petunia, or genes exogenous to plents

(e.g., genes derived from animals, inscots, bacteria, and tung), where the expression of products of such a gene are desired in microspores. A prefereble example of such a heterologous gene in the present Invention is a gene which encodes a cytotoxic gene product and whose expression inhibits the development of pollen. As especific example of such a gene, the barnese gene especific example of such a gene, the barnese gene (Beals, T. P. and Goldberg, R. B. P. |lent Cell, 31527, 1997) is illustrated, but the present invention is not limited to this.

[0059]. "Plent arprission vector" refers to a nucleic cod sequence including an expression cassate end, in addition, various regulatory elements linked to the cassette in such a menner that the regulatory elements can be operated in host plant cells. Preferably, such a plant expression vector may include a terminator, a drug-resistant gene, and an enhancer. It is well known matter to those skilled in the art that the types of plant expression vectors and the types of regulatory elements used may be varied depending on host cells. Plant expression vectors used in the present invention may further have a T-DNA region. Tha T-DNA region increases the efficiency of gene introduction, particularly when Agrobacterium is used to transform a plant.

Terminator is a sequence which is located downstream of a region encoding a protein of a gene and which is involved in the termination of transcription when DNA is transcribed into mRINA, and the addition of applyA sequence. It is known that a terminator contributes to the stability of mRINA, and has an influence on the amount of gene expression. As examples of such a terminator, the terminator for the nopalline synthase gene (Tros.), and the SSS terminator of caulifiower mosaic virus (CaMV) are illustrated, but the terminator is for limited to these.

[0061] "Drug-resistant-gene" is desirably one that facilitates the selection of transformed plants. The neomyclin phosphotransferase ii (NPTII) gene for conferring kanemycin resistence, ond the hygromycin phosphotransferase gene for conferring hygromycin resistance may be preferably used, but the drug-resistant gene is not limited to these.

[0062] The plant expression vector of the present invention may be prepared using a gene recombinant technique well known to those skilled in the art. A plant expression vector is constructed, for example, preferably using pBI-type vectors or pU-type vectors, but the plant expression vector is not limited to these.

50 (Production of trensformed plant)

[0063] The thus-constructed expression cassette, or an expression vector including the expression cassette, may be introduced into desired plant cells using a known gene recombinant technique. The introduced expression cassette is present to be integrated into DNA in a plant cell. It should be noted that DNA in a plant cell includes not only chromosome but ilso DNA included

In various organelles included in a plant cell (e.g., a mitochondria, and a chloroplast).

[0064] In the present specification, the term "plant" includes any of monocotyledons and dicotyledons. Preferable plants are dicotyledons. Dicotyledons include any of Archichlamiidae and Sympetalidae. A preferable subclass is Sympetalidae. Sympetalidae includes any of Gentianales, Solanales, Lamiales, Callitrichales, Plantaginales, Campanulales, Scrophulariales, Rubiales, Dipsacales, and Asterales. A preferable order is Solanales. Solanales includes any of Solanaceae, Hydrophyllaceae, Polemoniaceae, Cuscutaceae, and Convolvulaceae. A preferable family is Solanaceae. Solanaceae includes Petunia, Datura, Nicotiana, Solanum, Lycopersicon, Capsicum, Physalis, Lyclum, and the like. Preferable genera are Petunia, Datura, and Nicotiana, and more preferably Petunia. The genus Petunia includes the following species: P. hybrida, P. axillarls. P. infleta, P. violacea, and the like. A preferable species is P. hybrida. "Plant" means phanerogamic plants and seed obtained from the plants unless other-

[0065] As exemples of "plant cells", cells in each tissue of plant orgens, such as flowers, leaves, roots, and the like, callus, and suspension cultured cells are Illus- 25 trated

[0066] For the purpose of introduction of a plant expression vector into e plant cell, a method well known to those skilled in the ert, such as an indirect method using Agrobacterium, and a method for directly introducing into cells, can be used. As such an indirect method using Agrobacterium, for example, a method of Nagel et al. (FEMS Microbiol. Lett., 67:325 (1990)) may be used. In this method, initially, Agrobacterium is transformed with a plant expression vector (e.g., by electroporation), and then the transformed Agrobacterium is introduced into a plant cell with a well-known method. such as a leaf disk method and the like. As a method for directly introducing a plant expression vector into a cell, an electroporetion method, particle gun, a calcium phosphate method, a polyethylene glycol method, and the like are illustrated. These methods are well known in the art. A method suitable for a plant to be transformed can be appropriately selected by those skilled in the art.

[0067] Cells Into which a plant expression vector has been introduced are screened for drug resistance, such as kanamycin resistance and the like, for example. A selected cell may be regenerated to a plant using a commonly used method

[0068] Whether or not an introduced plant expression 50 vector is operative in a regenerated plant can be confirmed with a technique well-known to those skilled in the art. For example, in the case where suppression of the expression of an endogenous gene is intended. such confirmation can be conducted by measuring the level of transcription with Northern blot analysis. In this manner, a desired transformed plant in which the expression of an endogenous transcription factor is sup-

pressed can be selected. For the purpose of the expression of a heterologous gene using a tissue-specific promoter, the expression of the heterologous gene can be confirmed usually by Northern blot analysis using RNA

extracted from a target tissue as a sample. The procedures of this analysis method are well known to those skilled in the art. [0069] Whether or not the expression of an endog-

enous transcription factor is suppressed in accordance with the method of the present invention so that pollen fertility is reduced can be confirmed, for example, by observing the form of the pollen of a plant, which is transformed by an expression vector including DNA encoding a transcription factor, with a microscope optionally after histochemically staining.

[0070] Whether or not a promoter is expressed specifically in a microspore in accordance with the method of the present invention can be confirmed by, for example, histochemically staining flower tissues including the anther in a plant transformed with en expression vector. In which a promoter is operatively linked to the GUS gene, by a commonly used method to detect the distribution of GUS activity.

### (Examples)

[0071] Hereinafter, the present Invention will be described based on examples. The scope of the present invention is not limited to the examples only. Restriction enzymes, plasmids, and the like used in the examples are available from commercial sources.

(Example 1: Construction of plant expression vector including polynucleotide encoding ZPT trenscription factors)

[0072] Out of the previously reported anther-specific ZF genes (Kobayashi et al., above), cDNAs of PEThy ZPT2-5 (ZPT2-5), PEThy ZPT3-1 (ZPT3-1), and PEThy ZPT4-1 (ZPT4-1) were each linked downstreem of the 35S promoter of the cauliflower mosaic virus to prepere a plant expression vector. This preparation will be specifically described below.

### (Example 1-1)

[0073] DNA fragments including the cauliflower mosaic virus 35S promoter (Hindill-Xbel fragment) and DNA fragments including the NOS terminator (Sad-EcoRI fragment) inplasmid pBI221 (purchased from CLONTECH Laboratories Inc.) were successively inserted into the multi-cloning site of plasmid pUCAP (van Engelen, F. A. et al., Transgenic Res., 4:288, 1995) to prepare pUCAP35S. A pBluescript vector including cD-NA of ZPT2-5 was cleaved at KpnI and SacI sites (either is a site within the vector), and inserted between KpnI and SacI sites of the above-described pUCAP35S. Further, this recombinant plasmid was cleaved with EcoRI

and Hindill, and a DNA fragment encoding ZPT2-5 was inserted between EoPli and Hindill sites of binary vector pBINPLUS (van Engelen, F. A. et al., above). As shown in Figure 4(a), the constructed ZPT2-5 gene comprises the SSS promoter region (P3SS: 0.9 kb) of cauliflower mosalic virus (CaMV), a polynucleotide (ZPT2-5; about 0.8 kb) encoding ZPT2-5 of the present invention, and the terminator region of nopaline synthase (Tnos: 0.3 kb). Pnos in Figure 4 indicates the promoter region of nopaline synthase, and NPTII indicates the necomicin phosphotrasterizes all gene.

### (Example 1-2)

[0074] A pBlusecript vector including cDNA of ZPT3-1 to was cleaved at Kpn I side and Soal side (either Is a site within the vector), and inserted between KpnI and Sac lates of pULRAPSS. Further, this recombinant plasmid was cleaved with EcoRI and HindlII, and a DNA fragment encoding ZPT3-1 was introduced between EcoRI and HindlII side of binary vector pBINPLUS. As is apparent from Figure 4(b), the constructed ZPT3-1 gene comprises the SSS promoter region (PSSS; 0.9 kb) of cauliflower mosalc virus (CaWV), a polynucleotide (ZPT3-1; about 1.7 kb) encoding ZPT3-1 of the present 2s invention, and the terminator region (Tnos; 0.3 kb) of nopaline synthase.

### (Example 1-3)

[0075] A pBluescript vector including cDNA of ZPT4-1 was cleaved at Kprl site and Sacl site (lether is a site within the vector), and inserted between KpnI and Sacl site of the above-described pUCAPSS. Futher, this recombinant plasmid was cleaved with EcoRI and Hindill, and a DNA fragment encoding ZPT4-1 was inful did and a DNA fragment encoding ZPT4-1 vas inful sites of binary vector pBINPLUS. As is apparent from Figure 4(c), the constructed ZPT4-1 gene comprises the 35S promoter region (PSSS; 0.9 kb) of cauliflower mosaic virus (CaMV), a polynucicolide (ZPT4-1; about 2.0 kb) encoding ZPT4-1 of the present invention, and the terminator region (Toss; 0.3 kb) of nopaline synthase.

(Example 2: Isolation of ZPT3-1 and ZPT4-1 promoter 45 regions and linkage to GUS reporter gene)

[0076] cDNAs of ZPT3-1 and ZPT4-1 were used as probes to isolate corresponding genomic clones from the genome DNA library of Petuna. DNA Iragements 90 (promoter region; about 2.7 kb and about 3.6 kb) upstream of the transcription initiation site were subcloned. Each DNA Iragement was linked upstream of the GUS reporter gene and cloned into a binary vector. This preparation will be specifically described below.

### (Example 2-1)

[0077] cDNA of ZPT3-1 was labeled with [\alpha-32P] dCTP using a commonly used random priming method (Sambrook et al., above) to prepare a radiolabeled probe. With this probe, a genomic library of Petunia (Petunia hybrida var. Mitcheli) prepared within EMBL3 vector (manufactured by Stratagene) was screened. A genome DNA fragment (Pstl-Sacl) of about 2.7 kb including the upstream region of the gene from the resultant clone was subcloned at PstI-Saci site of pBiuescriptSK vector (pBS-ZPT3-1-PS), followed by sequencing (Figure 5). Next, this plasmid was used as a template to conduct PCR using a primer including a Sail recognition sequence (3'-TATGGAGCTCGTCGACAG TTGATGGT-TCATTTTTCTGGCTATTGTC-5'; SEQ ID NO: 11) and a commercially available M13-20 primer, so that Sall site was introduced immediately downstream of the initiation site of translation of the ZPT3-1 protein (base position: 2661). Thereafter, a DNA fragment cleaved with Psti and Sall was Inserted upstream of the GUS coding region of pUCAPGUSNT (pUCAP-ZPT3-1-GUSNT). Therefore, the ZPT3-1 gene was connected to the GUS coding region in frame at a region near the N terminus of the coding region of the ZPT3-1 gene. Further, a DNA fragment obtained by cleaving pUCAP-ZPT3-1-GUSNT with AscI and Paci (including the ZPT3-1 promoter, the GUS coding region and the NOS terminator) was inserted Into pBINPLUS vector to obtain pBIN-ZPT3-1-GUS (Figure 7(a)).

### (Example 2-2)

[0078] As to ZPT4-1, similarly, genomic DNA was isolated, and a DNA fragment (EcoRi-EcoRI) of about 3.6 kb including an upstream region of the ZPT4-1 gene was subcloned at the EcoRI-EcoRI site of pBluescriptSK vector (pBS-ZPT4-1-EE), followed by sequencing (Figure 6). This plasmid was used as a template to conduct PCR using a primer including a BamHI recognition sequence (3'-CATGGATATAGGATCCTATATC-5'; SEQ ID NO: 12) and M13-20 primer, so that BamHI site was introduced Immediately downstream of the initiation site of translation of the ZPT4-1 protein (base position: 3641). Thereafter, a DNA fragment cleaved with EcoRI and BamHI was Inserted upstream of the GUS coding region of pUCAPGUSNT (pUCAP-ZPT4-1-GUSNT). Therefore, the ZPT4-1 gene was connected to the GUS coding region in frame at a region near the N terminus of the coding region of the ZPT4-1 gene, Further, a DNA fragment (AscI-PacI) was inserted into pBINPLUS vector to obtain pBIN-ZPT4-1-GUS in a manner similar to that described above (Figure 7(b)).

55 (Example 3: Introduction of each fusion gene into Petunia cells)

[0079] Each of the above-described expression vec-

tors was introduced via Agrobacterium into Petunia (Petunia hybrida var. Mitchell) with the following procedures.

- (1) Agrobacterium tumefaciens LBA4404 strain (purchased from CLONTECH Laboratories line.) was cultured at 28°C in L medium containing 250mg/ml of streptomycin and 50mg/ml of infampicin. Cell suspension was prepared in accordance with the method of Nagel et al. (1980) (above). The plasmid vector constructed in Examples 1 and 2 were introduced into the above-described strain by electroporation.
- (2) A polynucleotide encoding each fusion gene was introduced into Petunia cells using the following method: the Agrobacterium tumefaciens LBA4404 strain obtained in the above-described (1) was shake-cultured (28°C, 200 rpm) in YEB medium (DNA Cioning, Vol. 2, page 78, Glover D. M. Ed., IRL Press, 1985). The resultant culture was diluted with sterilized water by a factor of 20, and cocultured with leaf pieces of Petunia (Petunia hybrida var. Mitcheil). After 2 to 3 days, the above-described bacterium was removed in medium containing antiblotics. The Petunia cells were subcultured with selection medium every two weeks. The transformed Petunia cells were selected based on the presence or absence of kanamycin resistance due to the expression of the NPTII gene derived from 30 pBINPLUS which had been introduced along with the above-described five fusion genes. The selected cells were induced into callus with a commonly used method. The callus was redifferentiated into a plant (Jorgensen R. A. et al., Plant Mol. Biol., 31: 35 957, 1996).

(Example 4: Phenotype of transformed Petunia into which ZPT genes are Introduced)

[0080] The transformants obtained by introducing the vector of Example 1 were used to observe change in the form of pollon in association with the control of the expression of ZPT2-5, ZPT3-1 and ZPT4-1, so that the influence of the introduced CDNA of these ZPT genes on plants were studied. This study will be described below in detail.

(Example 4-1)

[0081] From transformants (14 individuals) into which cDNA of ZPT2-5 had been introduced under the control of a 35S promoter, individuals (3 individuals) in which gene expression was suppressed by cosuppression were selected by Northern blot analysis (note that over 55 expression of the ZPT2-5 gene introduced was observed in four individuals out of the 14 individuals). The conditions of the Northern blot analysis were the follow-

ing: hybridization was conducted in a solution containing '7% SDS, 60% formamide, 5x-SSC, 2% blocking reagent (manufactured by Boehringer Mannheim), 50 mM sodium phosphate buffer (plr 1.0), 0.1% sodium lauryl sarcosine, 50 µpml of yeast IRNA, and <sup>292</sup>-labeled probe DNA (1x10° cmp) at 68°C for 16 hours, followed by washing with 2x SSC/0.1% SDS at 88°C for 30 minutes. [0082] In the above-described three cosuppression transformants, the following phenotypes were observed (Floure 8).

(Figure 8).
[0083] In the meiosis process which occurs immediately before the tetrad stage, in the case of normal (wid type) Petunia, chromatin is condensed into thin thread-like structures (prophase I: leptotene), and synapsis of homologous chromosomes occurs (prophase I: zygotene). Thereafter, in metaphase I, chromosome tetrad sign along the equatorial plane of the cells, and thereafter the homologous chromosomes are equally separated to the opposite poles of the cells by the spindle apparatus. In the transformant having cosuppression of the 2PT2-5 gene, the separation of the chromosomes to the poles proceeded while chromosome tetrad did not align along the equatorial plane in metaphase I. The division of the chromosomes to the poles was significantly unbalanced.

[0084] In the normal process of melosis, after the above-described first separation of the chromosomes. second separation of the chromosomes forms four haploid groups. Thereafter, separation of cytopiasm occurs. In the case of the above-described transformant having cosuppression, separation of cytoplasm and cell divislon occurred immediately after the first separation of chromosomes. This unbalanced cell division occurred not only at a single time but also further repeated at least two times, so that 8 microspore cells were formed at the most. Due to the unbalanced separation of chromosomes, the number of chromosomes included in the microspore cells was unequal and, in addition, the size of the cells was significantly unequal. As a result, during the stage corresponding to the tetrad stage of normal Petunia, a more number of microspores (8 or less) than normal were formed in these transformants (Figure 8(f): a photograph of polien cells of the ZPT2-5 cosuppression transformant in the bud having a size of 6 mm. [0085] Further, Figure 9(b); see a photograph of pol-

len cells of the transformant in the tetrad stage).

[0086] In the cosuppression transformants, a part of
the microspores (10-20%) still continued to develop, but
most microspore cells burst before a caliose slear-onselection of the microspore was degraded. In this stage,
the microspores which did not burst and survived were
in the abnormal form of substantially a haxhadron,
which was clearly different from the tetrahedron form of
normal microspores. Thereafter, the abnormal-form micorpores became binuclear due to seemingly normal
mitosis to form pollen grains. However, most of these
pollen grains lost fertility. Specifically, when the pollen
grains of these transformants were pieced on the pistil

of normal Petunia, no or few seeds were formed from pollen of the three strains exhibiting cosuppression (10% or less, i.e., the number of seeds produced by one Petunia is 10% of control as the average of about 10 (lowers). For pollen from three transformant strains without cosuppression, normal seed formation was confirmed similar to wild two control plants.

[0087] The above-described cosuppression transformat also exhibited abnormality in formation of fomale gametophyte, and formale fertility was reduced to 25-35% of that of normal individuals. Specifically, the development of an ovule (female gametophyte) was seemingly normal, but when wild type pollen was used for pollination, the majority of ovules could not be fertilized and even tertilized ovules exhibited abnormality in the subsequent developement, so that most ovules aborted. In this case, the transformants without cosuppression formed normal female gametophytes similar to wild type control plants.

### (Example 4-2)

[0088] CDNA of ZPT3-1 was introduced under the control of the 35 promoter. As a result, a trait change similar to when the ZPT2-5 gene was introduced appared in their endividuals out of 15 find/dutals (Figure 9). Specifically, in these transformants, substantially the same abnormally as that of ZPT2-5 were observed in the process of melosis. The number of cells which developed up to the microspore stake was very small, and 30 surviving microspores exhibited morphological abnormality (hexabedron). Further, matured pollen grains lost fertility. However, unlike ZPT2-5, the female fertility of these Ind/dutals was not affects.

[0089] Gene expression was analyzed with the Northern blot method under the same conditions as those in Example 4-1. As a result, in individuals into which the ZPT3-1 gene was introduced, gene expression was suppressed both for ZPT3-1 and ZPT4-1. Both genes share a high level of structural similarity. Specifically, the homology of the base sequence in the entire coding region is 37%. When the second ZF region of ZPT3-1 and the third ZF region of ZPT4-1, and the third ZF region of ZPT3-1 and the fourth ZF region of ZPT4-1, including neighboring sequences, are respectively compared with each other at the base sequence level in such a manner that the homology value is maximized, the average of the homology is 86% (the comparison of the sequences was conducted using the Clustal V program). Therefore, it is highly likely that the above-described expression 50 suppressing phenomenon is caused by the introduction of one gene leading to the suppression of the expression of two genes (cosuppression). This suggests that the functions of these two genes overlap, and is consistent in that by the introduction of either gene, a common 55 change in a phenotype could be observed.

### (Example 4-3)

[0090] cDNA of ZPT4-1 was introduced under the control of the 35S promoter. As a result, a trait change similar to when the ZPT2-5 gene was introduced appeared in two individuals out of 1s individuals. Specifically, in these transforments, substantially the same abnormality as that of ZPT2-5 were observed in the process of melosis. The number of cells which developed up to the microspores stage was very small, and surviving microspores exhibited morphological abnormality (hexahedron). Further, nest matured pellen grains lost fertility. However, similar to ZPT3-1, the female fertility of these individuals was not affected. For the above-described resons, in this example, it is also highly likely that gene expression was suppressed for both ZPT3-1 and ZPT4-1 (cosuppression).

[0091] As described above, by introducing a gene encoding ZPT2-5, ZPT3-1 or ZPT4-1, the development of pollen can be inhibited and the fertility can be eliminated with excellent efficiency (995 or more for ZPT3-1, and 90% or more for ZPT2-5 and ZPT4-1). The introduction of these genes may be useful for a selective trail transformation technique since the effects of the genes are specific to polen (pollen and female gamesophy in in the case of ZPT2-5) and the other traits of plants are not affected.

(Example 5: Tissue specificity of promoter activity of ZPT3-1 and ZPT4-1)

[0092] The tissue-specific promoter activity of the above-described DNA fragments was detacted by histochemical staining with GUS activity using the transformants obtained by introducing the vector in Example 2. This will be described below in detail.

### (Example 5-1)

40 [0093] The flowers of the transformants obtained by introducing a fusion gene of the upstream region of the ZPT3-1 gene with GUS were used to study the distribution of GUS activity using X-GUS as a substrate [Guspher, S. R. Ed., GUS protocols: using the GUS gene 45 as a reporter of gene expression, Academic Press, Inc., pp. 103-114, 1992). As a result, GUS activity was detected specifically in microspores in the uninucleate state (Floures 10fe) through (cf.).

### 50 (Example 5-2)

[0094] The flowers of the transformants obtained by introducing a fusion gene of the upstream region of the ZPT4-1 gene with GUS were used to study the distribution of GUS activity in a manner similar to that described above. As a result, GUS activity was observed specifically in microspores and the dehiscence tissue of anterior from the uninucleate stage to the binucleate stage.

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(Figure 10(d) through (f); the dehiscence tissue of anthere was Indicated by an arrow in Figure 10(g) and (f)). [0095] As described above, the promoters for the 2F73-1 and 2F74-1 genes exhibit activity specifically in microspores in the uninucleate stage (2F73-1) and microspores from the uninucleate stage (2F73-1) and microspores from the uninucleate stage to the binucleate stage (2F74-1), respectively. The promoter for the 2F74-1 gene also exhibits activity specifically in the dehiscence tissue of anthers from the uninucleate stage to the binucleate stage.

[0096] Microspores are precursor cells which will be subsequently matured to form pollen grains. Therefore, those promoters are useful as a tool for detailed research on the development of pollen. Further, these promoters or active fragments thereof can be used to cause 15 a cytotoxic gene or the like to be expressed specifically in microspores to abort pollen cells or eliminate the functions thereof, whereby the development of pollen can be directly and efficiently controlled.

### INDUSTRIAL APPLICABILITY

[0097] The method of the present invention utilizing DNA encoding transcription factors derived from the ZPT2-5, ZPT3-1 and ZPT4-1 genes, and promoters derived from the ZPT3-1 and ZPT4-1 genes is useful as a technique for selectively modifying the trait of a plant using a genetic engineering method, particularly a technique for conferring male sterified.

### Claims

 A method for producing a male sterile plant, comprising the steps of:

providing a plant expression cassette including: a nucleic acid being any of (), DNA having a sequence from position 1 to position 777 of a base sequence indicated by SEO IO NO: 1, (ii) 40 DNA hybridizing the DNA having the base sequence ellipated to the providing and encoding a transcription factor controlling the development of polion, and (iii) a DNA fragment of (i) or (II); and a promoter operatively linked to the nucleic acid:

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions; introducing the expression cassette into the

plant cells; regenerating the plant cells, into which the expression cassette has been introduced, to 55

screening the regenerated plants for one in which the nucleic acid is expressed so that ex-

plants: and

pression of the endogenous transcription factor is suppressed,

wherein the DNA of (ii) does not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of a base sequence indicated by SEQ ID NO: 13 under stringent conditions and encoding a transcription factor controlling the development of pollen.

A method for producing a male sterile plant, comprising the steps of:

providing a plant expression cassette including: a nucleic acid being any of (f) DNA having a sequence from position 1 to position 1840 of a base sequence from position 1 to position 1840 of a base sequence indicated by SEQ ID NC: 3, (iii) DNA hybridizing the DNA having the D

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions;

introducing the expression cassette into the plant cells;

regenerating the plant cells, into which the expression cassette has been introduced, to plants; and

screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed.

wherein the DNA of (ii') does not include (iv) DNA hybridzing DNA having a sequence from position 1 to position 1886 of a base sequence indicated by SEQ IO NO: 13 under stringent conditions and encoding a transcription factor controlling the development of pollen.

A method for producing a male sterile plant, comprising the steps of:

providing a plant expression cassette including: a runciets acid being any of ("DNA having a sequence from position 1 to position 1 ske of a base sequence indicated by SEO (IDNO.5, 6)(") DNA hybridizing the DNA having the base set of the plant of the p

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nuclelc acid under stringent conditions:

introducing the expression cassette into the plant cells;

regenerating the plant cells, into which the expression cassette has been introduced, to plants; and

screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed.

wherein the DNA of (ii'') does not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of a seas sequence indicated by SEQ ID NO: 13 under stringent conditions and encoding a transcription factor controlling the 20 development of pollen.

- A method according to any of claims 1 through 3, wherein the nucleic acid is linked in a forward direction with respect to the promoter, and may be transcribed in a sense direction in cells of the plant.
- A method according to any of claims 1 through 3, wherein the nucleic acid is linked in a reverse direction with respect to the promoter, and is transcribed in a antisense direction in cells of the plant.
- A method according to any of claims 1 through 3, wherein the plant is dicotyledon.
- A method according to claim 6, wherein the plant is of the family Solanaceae.
- A method according to claim 7, wherein the plant is of the genus Petunia.
- A method according to any of claims 1 through 3, wherein the expression cassette is incorporated into a plant expression vector.
- A maie sterile plant produced by a method according to any of claims 1 through 9.
- 11. A method for conferring male sterility to a plant, comprising the steps of:

providing a plant expression cassette including: a nucleic acid being any of (i) DNA having a sequence from position 1 to position 777 of a base sequence indicated by SEG ID NO: 1, (ii) DNA hybridizing the DNA having the base sequence (i) under stringent conditions and encoding a transcription factor controlling the de-

velopment of pollen, and (iii) a DNA fragment of (i) or (ii); and a promoter operatively linked to the nucleic acid;

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions:

introducing the expression cassette into the

regenerating the plant cells, into which the expression cassette has been introduced, to plants; and

screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed,

wherein the DNA of (ii) does not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of a base sequence indicated by SEQ ID NO: 13 under stringent conditions and encoding a transcription factor controlling the development of pollen.

A method for conferring male sterility to a plant, comprising the steps of:

providing a plant expression caseata including: a nucleic acid being any of (7) DNA having a sequence from position 1 to position 1846 of a base sequence indicated by SEC (10 NO: 3, (1)) DNA hydriding the DNA having the base sequence (1) under stripgent conditions and encoding a transcription factor controlling the development of poline, and (iii) a DNA tragment of (7) or (iii); and a promoter operatively linked to the nucleic acid.

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions:

introducing the expression cassette into the plant cells;

regenerating the plant cells, into which the expression cassette has been introduced, to plants; and

screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed,

wherein the DNA of (ii') does not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of a base sequence indicated by SEQ ID NO: 13 under stringent conditions and encoding a transcription factor controlling the

development of pollen.

13. A method for conferring male sterility to a plant, comprising the steps of:

providing a plant expression cassette including: a nucleic acid being any of (") DNA having a sequence from position 1 to position 1948 of a base sequence indicated by SEGI DNC:5, (") DNA hybridzing the DNA having the base sequence (") under stringent conditions and encoding a transcription factor controlling the development of polien, and (iii) = DNA fragment of (") or (ii'); and a promoter operatively linked to the nucleic acid;

providing plant cells having an endogenous transcription factor controlling the development of pollen, wherein a gene encoding the endogenous transcription factor hybridizes the nucleic acid under stringent conditions;

introducing the expression cassette into the plant cells;

regenerating the plant cells, into which the expression cassette has been introduced, to plants; and

screening the regenerated plants for one in which the nucleic acid is expressed so that expression of the endogenous transcription factor is suppressed,

wherein the DNA of (III') does not include (iv) DNA hybridizing DNA having a sequence from position 1 to position 1886 of a base sequence indicated by SEO ID NO: 13 under stringent conditions and encoding a transcription factor controlling the 35 development of polien.

14. A method for producing a plant having a modified trait, comprising the steps of:

providing a plant expression cassetta including: a promoter including any of (a) DNA having a sequence from position 1 to position 2624 of a base sequence indicated by SEO (I) NO.7 and (b) DNA having a part of the sequence of (a) 45 and exhibiting promoter activity specifit to microspores; and a haterologous gene operatively linked to the promoter;

introducing the expression cassette into plant cells; and

regenerating the plant cells, into which the expression cassette has been introduced, to plants.

15. A method for producing a plant having a modified 55 trait, comprising the steps of:

providing a plant expression cassette including:

a promoter including any of (a\*) DNA having a sequence from position 1 to position 3831 of a base sequence indicated by SEQ ID NO: 8 and (b\*) DNA having a part of the sequence of (a\*) and exhibiting promoter activity specific to microspores and optionally the dehiscence tissue of an anther; and a heterologous gene operatively linked to the promoter; introducing the expression cassette into joint

cells; and regenerating the plant cells, into which the expression cassette has been introduced to

plants.

16. A method according to claim 14 or 15, wherein the

- trait is fertility, and the plant having a modified trait is a male sterile plant.
- 17. A method according to claim 14 or 15, wherein the trait is compatibility, and the plant having a modified trait is a self-incompatibile plant.
- A method according to claim 14 or 15, wherein the plant is dicotyledon.
- A method according to claim 18, wherein the plant is of the family Solanaceae.
- A method according to claim 19, wherein the plant
   is of the genus Petunia.
  - A method according to claim 14 or 15, wherein the expression cassette is incorporated into a plant expression vector.
  - A trait-modified plant produced by a method according to any of claims 14 through 21.
  - 23. A method for conferring male sterility to a plant, comprising the steps of:

providing a plant expression cassette including: a promoter including any of (a\*) DNA having a sequence from position 1 to position 2624 of a base sequence indicated by SEQ ID NO; 7 and (h\*) DNA having a part of the sequence of (a\*) and exhibiting promoter activity specific to microspores; and a heterologous gene operatively linked to the promoter,

introducing the expression cassette into plant cells; and

regenerating the plant cells, into which the expression cassette has been introduced, to plants.

24. A method for conferring male sterility to a plant, comprising the steps of:

providing a plant expression cassette including: a promoter including any of (a") DNA having a sequence from position 1 to position 3631 of a base sequence indicated by SEO ID NO: 8 and (b") DNA having a part of the sequence of (a") and exhibiting promoter activity specific to microspores and optionally the dehiscence tissue of an anther; and a heterologous gene operatively linked to the promoter;

introducing the expression cassette into plant 10 cells; and

regenerating the plant cells, into which the expression cassette has been introduced, to plants.

25. A promoter comprising DNA of the following (I') or (II'):

(l') DNA having a sequence from position 1 to position 2624 of a base sequence indicated by 20 SEQ ID NO: 7 and (il') DNA having a part of the sequence of (i') and exhibiting promoter activity specific to microspores.

A promoter, comprising DNA of the following (i") or 25 (II"):

(I") DNA having a sequence from position 1 to position 3631 of a base sequence indicated by SEQ ID NO: 8 and (ii") DNA having a part of 90 the sequence of (i") and exhibiting promoter activity specific to microspores and optionally the dehiscence itsisue of an enther.

27. A plant expression cassette useful for conferring male sterility to a plant, comprising a promoter according to claim 25 or 26 and a heterologous gene operatively linked to the promoter.

55

# FIG. 1

atca	aaaa	ca a	aati	ccti	t ti	Cac	accga	a aga	acag	ject	tagt	atti	ca a	ngaaa	ac	57
											gaa Glu					105
											ctc Leu					153
											agt Ser					201
							Arq				tet Ser 60					249
											tta Leu					297
										Lys	ogt					345
									Ile		Caa Gln			Gly		393
			Arq								Lys		Leu		got Ala	441
		Asp									gca Ala 140	Asn				489

# FIG. 1 (Continued)

att	ttg	tcc	ttg	gat	ttg	aac	ttg	acg	cca	ttg	gaa	aat	gac	tta	gag	537
Пe	Leu	Ser	Leu	Asp	Leu	Asn	Leu	Thr	Pro	Leu	Glu	Asn	Aap	Leu	Glu	
145					150					155					160	

ttt gat ttg oga eag agt aat act got oot atg gto gat tgo ttt tta 585 Pho Asp Leu Arg Lys Sex Asm Thr Ala Pro Met Val Asp Cys Pho Leu 165 170

tga ttgaacttto cgtttootta ttottttoto ttottottt ggatattgta 638 tttattoatt aattgtagga gggataggaa gtottatott gtgtattagt ectacenttt 658 gcagattgta gaacgattag tttgtaactt atcatgatac cogaantaca atactattta 758 tatgattatt atactacac

# FIG.2

accegatecaga aattecegaga tegacecaca egtecagaaa ettteettat tacaetttaa 60	<b>)</b> .
thatethet agreements themesens and a single	
ttatgttot agtgagtata ttagagagtg agaa atg gtg gac aat agc cag aaa <u>1</u> Met Val Asp Asn Ser Gln Ivs	L5
1 5	
aat gaa oo a toa act git ata cac tat tgi aga gia tgi aaa agg gga 16	2
Asn Glu Pro Ser Thr Val Ile His Tyr Cys Arg Val Cys Lys Arg Gly	
10 15 20	
itt aat agt got gga got ott ggt ggg oac atg aga tot oat gga gtg 21	1
The Asm Ser Ala Cly Ala Leu Cly Cly His Met Arg Ser His Cly Val	
25 30 35	
gga gat cat aat aaa aac tat ggt gaa gat att aat gaa caa aga tat 25	_
aga gat cat aat aan aan tat ggt gan gat aft ant gan can aga tat 25 My Asp His Asn Lys Asn Tyr Gly Glu Asp Ile Asn Glu Gln Arg Tyr	9
40 45 50 55	
-5 50 . 55	
atg atc aac aac ttt aga aga gat aaa cca gag ggt caa aag cac tca 30	7
det Ile Asn Asn Phe Ary Ary Asp Lys Pro Glu Gly Gln Lys His Ser	•
60 65 70	
tat aat oft ogt got aat act aat aga tta tta ggo aat oga goa agt 35	5
Tyr Asn Leu Arg Ala Asn Thr Asn Arg Leu Leu Gly Asn Arg Ala Ser	
75 80 85	
was gat cgt gac aag aag too tog atg tgg oot ooc aat gat cgt ggg 40	3
Elu Asp Arg Asp Lys Lys Ser Ser Met Trp Pro Pro Asn Asp Arg Gly	
90 95 100	
asa tat goc cta gac gag act cta acc cta toa toa atg tog toa coa 45	
lys Tyr Ala Leu Asp Glu Thr Leu Thr Leu Ser Ser Met Ser Ser Pro	Ξ,
105 110 115	
gga toa toa gat ott gaa aga agt act aag oca tat gat goa aaa gaa 49	9
Bly Ser Ser Asp Leu Glu Arg Ser Thr Lys Pro Tyr Asp Ala Lys Glu	
L20 125 130 135	

FIG	?. <i>2</i>	(C	ontir	nued	1)											
														gea Gl.u 150		547
														tat Tyr		595
													Val	gaa Glu		643
												Ser		cac His		691
											Val			tgt Cys	tat Tyr 215	739
														aac Asn 230		787
														oct Pro		835
													Gln	tot Ser		883
												Val.		caa Gln		931
tcg	att	tat	cat	cqa	att	ttt	tca	tca	gga	Caa	acc	tta	gat.	aaa	cac	979

Ser Ile Cys His Ard Val Phe Ser Ser Gly Gln Ala Leu Gly Gly His

$FIG.\mathcal{Z}$ (Continued)	
ana agg tgt cac tgg cta tca tca agt ttg cca	
Lys Arg Cys His Trp Leu Ser Ser Ser Leu Pro	
300 305	
303	310
cca act ttt caa gaa atc caa tao cac acc caa	gaa caa gga tta tto 1075
Pro Thr Phe Gln Glu Lle Gln Tyr His Thr Glr	
315 320	325
aac aag coa atg ttt acc aac ttt gat caa cos	tta gat cta aac tto 112
Asn Lys Pro Met Phe Thr Asn Fhe Asp Gln Pro	Leu Asp Leu Asn Phe
330 335	340
oca gca caa cta ggc aat cca gct gaa ttt gag	
Pro Ala Gln Leu Gly Asn Pro Ala Glu Phe Glu	
345 350	355
oca ttt gaa cat gaa ggc cca aga agc tat ctc Pro Phe Glu His Glu Gly Pro Arg Ser Tyr Lea	
360 365 37	
	3,3
caa caa atc aat act aat tta cat caa aat gag	g ang tgo ana gat toa 126
Gin Gin Lie Asn Thr Asn Leu His Gin Asn Gli	u Lys Cys Lys Asp Ser
380 385	390
acy gag gat ttg aga agg gaa gaa aat tac aag	
Thr Glu Asp Leu Arg Arg Glu Glu Asn Tyr Lys	
395 400	405
ttg agt aac ctt aaa gat gtg aac ttg gat ggs	
Leu Ser Asn Leu Lys Asp Val Asn Leu Asp Gly	a ggo tot tot tgg tta 1363
410 415	420
	420
caa gta ggg att ggt cca acc cca gat ata gta	a goa act ctg taa 1408
Gin Val Gly Ile Gly Pro Thr Pro Asp Ile Val	
425 430	435
ggttagtaac acagtgatcg ttatgtcagc tacaagtate	
ocaacttata cataaactgt ttaacatatt tatacttto	g tattattgtt gtatcgaact 152

ttoactagtt acaatttgtg attogtocaa tooctaatat agtagcaaca gacctgtaag 1588 attagtatta tgcgattgtt ttgtcattct acaaaataaa atcgtataat at

# FIG.3

		_														
0000	cat	gca e	ttt	tttt	eg to	stati	cati	t cto	ctca	scta	aaaa	ctaga	att 1	tgeti	totta	t 60
agti	tct	gt d	cat	gtoto	at te	stcat	ttcat	t ac	ttgae	agta	gta	caata	ac a	aaga	aata	a 120
cati	tage															t 171
		M		sp C	ys I	le A		ln G	lu G	ln G			ln G	ln P	ro Va	a.
			1				5				3	LO				
ttt	aaq	cat	tat	tat	aga	att	tac	220	aaa	ant	***	ata	tet		aga	219
									Lys							219
15					20					25					30	
gat	cta	ggt	888	cat	atg	aga	gct	cat	gga	att	999	gat	gaa	gtt	gta	267
Ala	Leu	Gly	Gly		Met:	yra	Ala	His	СĨΆ		αъ	qaA	Gш		Val	
				35					40					45		
act	ata	cet	cent	cat	cont	can	ന്നു	acrit	gat	+	<b></b>	-		+++		315
									Asp							313
			50	_	_			55					60			
899	agt	gtt	aag	gaa	ggt	aat	888	égg	atg	tac	caa	tta	aga	aca	220	363
G13	Ser		Lys	Glu	GΙΆ	Asn		Arg	Met	Tyr	Glu		Arg	The	Asn	
		65					70					75				
act	aat	200	CRR	222	acc.	aat	acra		tgt	~~		+				411
									CAS							#TT
	80	_		-		85					90					
ttc	tot	tot	tgg	888	tct	ttt	ctt	gaa	cat	gga	aaa	tgt	ago	tca	gaa	459
		Ser	Tap	LVS			Leu	Glu	His	Gly	Lys	CVS	Ser	Ser		
95					100					105					110	
œt	ara.	~~~	man	+~+	++=	at-	+	+	œc							
									Pro							507
				115					120	GL.y		GLU.	GL.Y	125	, L	
tac	att	tat	gat	gga	aga	aaa,	gaa	aaa	gga	tac	gga	tgg	tot	azz	aga	555
									Glv							

r.	10	·. U	loc	mui	lucu	,											
	aag	agg	tca	tta	aga	aca	aaa	gta	gga	ggc	ctt	agt	act	tca	act	tat	603
	Lys	Arg	Ser	Leu	Arg	Thr	Lys	Val	Gly	Gly	Leu	Ser	Thr	Ser	Thr	Tyr	
			145					150					155				
	CBB	tca	agt	gag	gaa	gaa	gat	ctt	cta	ctt	gca.	aaa	tgc	ctt	ata	gat	651
	Gln	Ser	Ser	Glu	Glu	Glu	Asp	Leu	Leu	Leu	Ala	Lys	Cys	Leu	Πe	Asp	
		160					165					170					
	tta	gcc	aat	gca	agg	gtt	gat	aca	tca	ttg	gtt	gag	oca	gaa	gag	tot	699
	Leu	Ala	Asn	Ala	λrg	Val.	Asp	Thr	Ser	Leu	Val	Glu	Pro	Glu	Glu	Ser	
	175					180					185					190	
															tog		747
	Сув	Ala	Ser	Ala	Ser	Arg	ŒШ	Glu	Glu	Arg	Ala	Ala	Arg	Asn	Ser	Met	
					195					200					205		
															gaç		795
	Ala	Tyr	Gly		Thr	Pro	Leu	Val		The	Arg	Val	Pro	Phe	Asp	Asn	
				210					215					220	•		
															gct		843
	ГÃа	Ala		GLY	Ala	Ser	Ser		GLY	Leu	Phe	Glu			Ala	Cys	
			225					230					235				
															gca		891
	TAR	240	VAL	Me	ASI	ser	245	GIII	ALA	Leu	GIA			Arq	Ala	Ser	
		240					245					250	,				
	cac			att	974		++							1	ata		
															Leu		939
	255	272	Lys	,	272	260	Cyn	- X-	ALG	Au	265		vab	GIII	Lieu	270	
	200					200					200	•				2/0	
	crat	ato	++=	p++	-	mat.	<b></b>	era+	-+-		***				caa		007
															Gln		987
			1.00	110	275	برمم	ш.	лор	Val	280	116	ш	LUS	ASD	285		
					213					200					203		
	tte	ctr	cae	acri	tce	882	+~	atr	acc		+		24-		gaa	+	1025
															Glu		1035
		200	ш.	290	Jer	-ys	Jer	1-20	295	ъÃа	-	nys	1115			CAR	
				290					293					300			

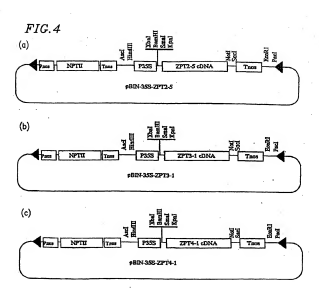
# FIG.3 (Continued)

tca	ata	tgc	cat	aga	gtt	tta	tcg	aca	gga	caa	gct	tta	ggt	ggt	cac	1083
Ser	Lle	Cys	His	Arg	Val	Phe	Ser	Thr	Gly	Gln	Ma	Leu	Gly	Gly	His	
		305					310					315				
aag	agg	tga	cac	tgg	atc	acc	tœ	aat	tco	œ	gat	tct	tcg	aaa	ttt	1131
Lys	Arg	Cys	His	Trp	Ile	Thr	Ser	Asn	Ser	Pro	Asp	Ser	Ser	Lys	Phe	
	320					325					330					
cat	ttc	aat	ggt	cat	gtg	gag	caa	att	aat	ota	aga	tca	aac	atg	cat	1179
His	Phe	Asn	GΙΥ	His	Val.	Glu	Gln	Пe	Asn	Leu	Arg	Ser	Asm	Met	His	
335					340					345					350	
888	tca	gat	gca	tta	gat	ott	aat	aac	ctt	œg	aca	cat	gaa	gac	atg	1227
Lys	Ser	Двр	Ala	Leu	Asp	Leu	maA	Asn	Leu	Pro	Thr	His	Glu	Asp	Met	
				355					360					365		•
tcg	cga	att	aga	cga	gac	000	ttt	aat	cca	tta	agc	ttc	gag	gtg	tca	1275
Ser	Arg	ملا	Arg	Arg	qaA	Pro	Phe	Asm	Pro	Leu	Ser	Phe	Glu	Val.	Ser	
			370					375					380			
	-			_					-	_	_	<u>~</u>			_	1323
Thr	qaA		His	Leu	Gln	Tyr		Trp	Ser	Cys	Ala	Pro	Lys	yeu	<b>qaA</b>	
		385					390					395				
		-										gat	-		-	1371
Asp		-	Asm	TYT	Tyr		GIU	Glu	He	Lys		Asp	Ser	Asn	Ala	
	400					405					410	•				
			_							-		caa		_	_	1419
		GIĀ	гåа	ıyr		ше	ASID	AST	GIĀ			Gln	AST	Val		
415					420					425	•				430	
-	-	-	-	-	_		_	_		_	_	cta	-	-		1467
Asp	'Asp	GIU	AIA	_		rÃa	Leu	LYP		ALA	TAB	Leu	ser			
				435					440					445		
										_		tgg				1515
TAB	AST	Met			Asn	Ser	Asp		PTO	Ala	His	Trp			val	
			450					455					460	)		

# FIG. 3 (Continued)

ggg att ggt toa act aca gaa gta ggg gct gat toa taa gtaactatat 1564 Gly Ile Gly Ser Thr Thr Glu Val Gly Ala Asp Ser 465 470 475

gcagttatto cittgottaa titottitit tiotgicaco ogagtatata titatatgoa 1624 aataitgidaa titataacito accaaacaga tagiaacigi tiggigatgo aaataoigit 1684 aataitigida citcoottiti tittgicoott tiotigiaat igatacacaa totigiaatii 1744 tittgiacit toaatitoit gagoigidaat titoagigia atacagaaci cagaatatgi 1804 tattottigoa ataigaagit tagiatagoaa cagicaaaca ogatagiag aagiggitoig 1864 taatoootoo cactagitaa aagitgggat tagitaacoo acagitagi gggotgaatii 1924 tgaagtaaac ataigcagit atto



## FIG.5

ctgcaggcag caacattagg agattttcca gcaccaatot ccctatgtgc tataacttca 60 cttataggca tggtattgac tggaattgta caattgatac aacaagggto gttggagatt 120 quattoccco tottaagcat cogtgactta ataqqctact cogttattoot aattcatcaa 180 atateeetga aatteteaca ttaattatgt taatacagaa attetgagtt agatttgaet 240 tacatacqtt gatagootaa ataatttgta toataotaac gtttttttaa cotoatactt 300 tatattaact ttgaggtttg totaattttt tgtggttatc ataggcaggt atagttagtg 360 gagcatgtgt aagtttcaat aattgggcaa tgaagaaaag agggccagtc ttagtttcog 420 tattagtes tottogaact gtgataactg tegtacttte toctateace ttgaagtaca 480 caattactat gogaagutaa aacettatoo attiteacti ggatetaget tatatacagt 540 gtaaggaast tittacaata tittccaagt aactittaaa gacgattatc aataatcatc 600 ttttacttaa cotgatagtg taaatatatt ttttcacact tacaattact ttagttottt 660 tteagttgca teaaaattea aactteaaat gaottaactt etttttgcag eettggtggt 720 atotttetea totttaeggg tetgtattte gtgttatggg etaaaaggaa egaaggattt 780 ctanataata ccaactoctc agaaagtgag taccatottg agaagcotgt tttocattan 840 atticittit attoccaatt gtaatatgta gttagtitot atatacaact agaatccaac 900 atagagaaga gagagggaga gottgtttgt accasataga taacatgtat gttgatttaa 960 gtatoccata ttggtactgg aagtanactg ttaatgttgc ctgcgattca attgtccagt 1020 cottootota greagacagt gittasatate ecacatogia taaaaaatog attoototot 1080 cottatatgg tatttgacaa tootcacatt ttgagctaaa atttgggttg agttaatgca 1140 attytoxatt tottatoxat gtatttaato taggottgga gotaaaaata caaagcaaaa 1200 gagaagagag aasaagaaga aagaagagt attatoatag ttgatatttg aasaaatgca 1260 agttocaato ctagtaatat cttttatttt gcagtagcat gacggaatat gggaatcaac 1320 atgtagotgc ttttctggct ctatctaago coctottett ttaccatagt tttgttttttc 1380 attoactttt ggaagoagoa aggytagatt tagaocacaa atatgoaaat gtttttttt 1440 ttttttttt tgtaaagtot tagacotata tggagtataa cotttgggaa aggggattga 1500 atcastcatc atsatgtcac satcatgtag tactacattt tttottctto satttgagot 1560 actaqtttga cattteccaa gtaaattatg cttcaacact aquattetet tgtttatatt 1620 atotoattga agotatgott taaototott oottgagtgg attaaottga aaaagtaggo 1680 anaganatti atgagagtto tgatatogat atcatagagg acacanantt naganantgc 1740 gassagactt atscccaaca sagasaatat gascactagt atcgatcacc acccagattt 1800 acaatttaat gtactggtgt toaattttgt gottgcatcg actatttcac cgaatattta 1860 ttottattta taasaatato gaataactat gaccatcaaa gtttagoosa ataasatata 1920 aaaaagtato tatatoaota tagtaaactt tgtatttatt ggaattgaac toacacttot 1980 tecattacta ggtcaaatec cagaaggcat attataagtt tttgtttcaa agoctecaaa 2040 oxaaqtacac tcattttctt tttgaagaaa gcgaqttcat ttqtaggcta cgtgaatata 2100 actactttaa aatattgett tgtttegaat ttgccatgag ttactacatt cacacaaaat 2160 tottaatgog actcagagtg tgtgttttaa ttttctttta gagtgtttgt acttctatat 2220 gaqqqtcact agtaaagtag tccactaata ttacaaattc ttacattacg tacaatgtga 2280

# FIG. 5 (Continued)

ttttatgtoa gtagatttga ctgaatgcta taactacgag agttagaaat agtotttgcc 2340
aacxacatta taaactgaco ctocacttgt cataacaaaa totuttgtte toatocacaa 2400
otaactttaa ctagaaacta gyacttooot cacttatget acaaaaato ttataacctac 2460
accacacact ttagtactgt tcactaacta attotttatt tataccaaco ctggottgga 2520
gtgtagcaaa aaaatgtaca ctactocaaa gtaaacacta ttottigaa otttoottgt 2580
tgcacttaa tttatgttot cgtgagtata ttagagagtg agaaatggtg gacaatagcc 2640
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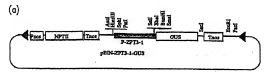
# FIG. 6

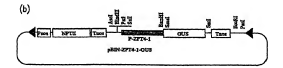
gaattoacca ccacqaqtac ttattttqat qaqcatqqca ttatttttca aacttottot 60 gotggaacac cacagcagaa cgggaaagtt gagcgaaaac ataaacatat tttgaatgtt 120 getegageae ttaggtttea agegeattte eeaattgagt tttggggtga gtgtgttttg 180 atggogtgot atttgatcaa gogaaccoto toatoggtot tacacagaaa aaaatgocat 240 atgatgtott ttttggtgta acaccgaact acgagcattt gaangtgttt ggatctctat 300 gotatggtca caagcatggg tgcttgggag ataagtttga aagtaggagt cgtotgtgtg 360 tittiatteg atacccatat gggaagaaat categaagtt atatgatite gacaccaaaa 420 satattttgt gtcgcgggca octagoscog aagcactaag catoogsacc caacgttatt 480 tragoctatg agagtgattg aagatgarta tggtattgaa gtgagggggt agtgaractg 540 ttttgacaca aaaaccgaac aaggagagga tacagutoga ggacgtgata attgacactc 600 caaqtttqqc tacagagact aatgtcatgg aagtagaaaa cocggtcacc ggtgtcatgt 660 cogatcaatt gaangotgaa gotgtgggag aagagttggg togagaaaac taattaggaa 720 ggagaatgto toottogtga tttttcactg gtotgtogaa aggttagtca ccaggtttca 780 acctgtgtcc acatgtctga tttctcaccc gtgacacaac gagcctcagg tacgccttat 840 ectettacae actatettaa tteteaceet ttttetteea accateteae ttttetteea 900 gotattacgg agggtogtga atogacctot ttotgtotog ccataaagaa tgaaaaatog 960 agazagacta tgczatagga gtttczagca ttgczagata atazaacatc tatogttout 1020 tacttgccac ctgggaagaa agcgotcgga tgtcggtogg tgtataagat caaatataat 1080 tocgatggat cagtggtacg atacaaggca cgtttggtta gttttggaaa toatcaggto 1140 asaggcattg attatacgta gacatttgct ccagtcgcta aaatagtgac tttgaggaca 1200 tttettgeag tegetgeage taanaatteg gaattgeate aaategatet teataatgea 1260 tttgtacagg tgatottcat gazaaagtot atatgaagot gocaccaagg tatcagacta 1320 atggttacgg taatgtgtgt cgcctatgaa agtttttgta tggtttgaag caggcgtcga 1380 gatgttggtt cacgaagtta ttggccgatt tgaaaactta tocttttana caatcttatt 1440 oggattattg octttttaca ottogtaaag ggtocgtcac ottaagtgtg ttggtgtacg 1500 tggatgattt gattattggg gcaataatto ggaagctatt cgtotottta agttgtatot 1560 ctocacttgc tttcttatga aagatttggg catactaant tttttgggag atgaagtggc 1620 tagaggacct aaaggtattt tootatgtca atggaaatat goottggata taattggatt 1680 attaggaget egactggttg gaacttetat ggageagaat categtttgg etttggcaag 1740 tggccgatat attgatgatc tacatagata tatttgattg atgattgtag tgcttaatta 1800 aagactgato aattgtactg ttattaatta atctttgttt aggaggagca tgtgggctgg 1860 assatgatgt agcasacttt coatacaatg coatgattac tocaqqasat gasgtoctat 1920 ttaaacatgg ctttggctgt ggtgcatgct accaggtgca cttgaaattt gttttataaa 1980 aagagaaaca catgcatgaa ttttgagttt cacttcgcaa aatanatgaa atctttattt 2040 atattaatgc aatcgatttt caggtgttgt gcttacagaa tcaaaatcaa tactgctcag 2100 gaaatocaat aatagtaact nttacagatg agtgcccagg ggcatgcaat aatgatoctg 2160 ttoattttga ttttagtgga actgcttttg gagccttggc aaaacctggc caagctgaac 2220 aattgogtaa tgaaggaaga atocaaatta attacagaag gtgagttacg ttocacatga 2280

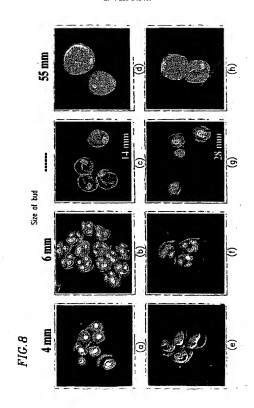
## FIG.6 (Continued)

caaatagaga aatcaataca aaatttocat ttacttagta acactctttc cttgttagta 2340 tgoctaaaaa agagtagtac acaacacaat taatgcacaa ttttgctaaa ccatgatatt 2400 gaatogtgca gagtggcatg cagttacaag gcasatatac aatttaaggt agacaaaggc 2460 tocaatoctg atticttggc agttgcagct gaggcagtta atggagatgg tgatctttct 2520 tttgtagaaa ttaaagcatc caattcgaat caatggette ceatgeaaca aatgtttggg 2580 goaacttgga gegttggeat caagecagae acaeagaaac eteetttete acttagaett 2640 actacagaat ttaagcaaac agtcattgoc caaaatgtca taccagtggg ttggcaacca 2700 agagcaattt acaaatcaaa tgtcaattte ccaccaage tttagtttaa tetttttacc 2760 cacaatagtg taaaaataat tataaggact acaaattaaa tagtotatgt toaacagtgc 2820 tatttaatta taataaggat tacaaattaa agtgaggatt ettotcaatg ataatgtcaa 2880 aagtttggga tgtcaaatot atttgtattt tttttcacat caatgatcaa tgaaagttat 2940 gottitagta ttttttaatt attaataatt tgttttcatg tatttcaata ataatattat 3000 otomangta antenatoma tattomanac tenceteman attiticatto tomotatatt 3060 tatgttottt titcagtoto aaacgoocaa attitgtacg aaaaaattgt toggataago 3120 gagaaacaot cataactgat aaaaacagaa tagtgaataa agaaaactaa atatatttac 3180 tettgatgag tecatgatgt gtaagtatta tottetgeeg tecaatttgg ttgtttgaca 3240 ocactagtgt tettaataan aagtttgtga aaaantaago tottoactoo ottaggoott 3300 actototoct tocactteto ataotoacto ttoacttoca etcacactoc tatttttete 3360 tttacotota aactotocto cacaaaccao taottcaaot aaaaaotagg actaattttt 3420 ttotcaccgt acaagtccac aacaacttot agtacaagaa caaacaaact ctcgttgtgc 3480 coolegates catgoatgea categories caattititt agtotottea tioteteaac 3540 taaaactaga tttgcttott atagtttott gtocatgtot ottotcatto atacttgaag 3600 tagtacaata acaaqaaat aacatttagc catggattgt atagatcaag aacaacaaca 3660 acaecaeca gttttteego attettgteg egtttgceeg eeeggttttg tgtgtgggeg 3720 agototaggt gggcatatga gagotoatgg aattggggat gaagttgtaa ctatggatga 3780 tgatgatcaa gcaagtgatt gggaagataa gtttggaggg agtgttaagg aaggtaataa 3840 aaggatgtac caattaagaa caaaccotaa taggcaaaaa agcaatagag titigtgagaa 3900 ttgtgggaaa gaattootgo agoooggggg atocactagt totagagong ngcgcacogo 3960 sytogagete cagettitet teeetttace teaggettaa tt 4002

# FIG.7







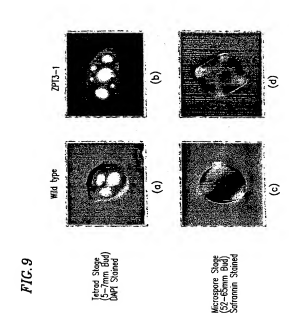
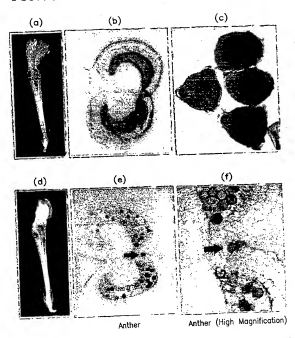


FIG. 10



	INTERNATIONAL SEARCH REPORT		International appl	ication No.
			PCT/J	P99/06467
A. CLASS Int.	IFICATION OF SUBJECT MATTER Cl <sup>7</sup> A01H5/00, C12N15/82			
According to	International Patent Classification (IPC) or to both nations	d classification or	M IPC	
	SEARCHED			
Minimum de Int .	currentation searched (classification system followed by c C1 A01HS/00, C12N15/82	lassification symb	ools)	
Jits	on searched other than minimum documentation to the extensive Shinan Koho 1922-1996 i Jitsuyo Shinan Koho 1971-2000	Toroku Jits	uyo Shinan K	in the fields searched Joho 1994-2000 Joho 1996-2000
Electronic di DIAL DDBJ	na base consulted during the international search (name of OG (BIOSIS)	data base and, wh	ere practicable, sea	urch terms used)
c. pocui	MENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where approp	riste, of the relev	ant passages	Relevant to claim No
A	The Plant Journal 13 [4] (1998) pp.571-576			1-27
A	Plant Molecular Biology 39 (1998), pp.1073-1078			1-27
A	Shivanna & Sawhney, Pollen biotechnology for crop production and improvement (1997) Cambridge University Press pp.237-257		,	1-27
Y A	Genes & Development, 5[3](1991) pp.496-507 pp.496-507			14-24 25-27
Y A	WO, 95/25787, A1 (RUTGERS UNIVERS 28 September, 1995 (28.09.95), Full text; Figs. 1 to 8 Full text; Figs. 1 to 8 & JP, 9-510615, A	ITY),	00	14-24 25-27
Further	documents are listed in the continuation of Box C.	See patent fami	ily annex.	
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m PCT/IS	A/210 (second sheet) (July 1992)			